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## Amendments to the Claims:

JC17 Rec'd PCT/PTO 23 JUN 2005

This listing of claims will replace all prior versions, and listings, of claims in the application:

## **Listing of Claims:**

- 1. (Original) A method for producing a carrier for the determination of analytes, comprising the steps:
  - (a) providing a carrier,
  - (b) passing liquid with building blocks for synthesizing polymeric receptors over the carrier,
  - (c) site- or/and time-specifically immobilizing the receptor building blocks on respective predetermined zones on the carrier and
  - (d) repeating steps (b) and (c) until the desired receptors have been synthesized on the respective predetermined zones,

characterized in that hapten groups are applied to the carrier before, during or/and after the synthesis of the receptors.

- 2. (Original) A method for the quality control of receptor syntheses on a carrier, comprising the steps;
  - (a) providing a carrier,
  - (b) applying in planar fashion hapten groups to the carrier surface,
  - (c) carrying out a receptor synthesis on the carrier,
  - (d) contacting with a hapten detection reagent which permits detection of hapten groups,

- (e) evaluating the hapten group detection on the carrier and
- (f) correlating the result of the evaluation with the quality or/and efficiency of the receptorsynthesis.
- (Original) A method for the quality control of receptor syntheses,
   comprising the steps:
  - (a) providing a carrier,
  - (b) carrying out a receptor synthesis on the carrier, with hapten groups being incorporated during the synthesis into the receptor molecules at predeterminedpositions,
  - (c) contacting with a hapten detection reagent which permits detection of hapten groups,
  - (d) evaluating the hapten group detection on the carrier and
  - (e) correlating the result of the evaluation with the quality or/and efficiency of the receptorsynthesis.
- 4. (Currently Amended) The method as claimed in any of claims 1 to 3 claim 1, characterized in that a microfluidic carrier with channels, preferably with closed channels, in which predetermined zones with immobilized receptors are produced is used.
- 5. (Currently Amended) The method as claimed in any of claims 1 to 4

<u>claim 1</u>, characterized in that the receptors are selected from biopolymers such as, for example, nucleic acids, nucleic acid analogs, proteins, peptides and carbohydrates.

- 6. (Currently Amended) The method as claimed in any of claims 1 to 5 claim 1, characterized in that the receptors are selected from nucleic acids and nucleic acid analogs.
- 7. (Currently Amended) The method as claimed in any of claims 1 to 6

  claim 1, characterized in that a carrier is produced with aplurality of, preferably with at least 50 and particularly preferably with at least 100, different receptor zones.
- 8. (Currently Amended) The method as claimed in any of claims 1 to 7

  <u>claim 1</u>, characterized in that the hapten groups are selected from organic molecules having a molecular weight of up to 2,000, which are recognized by a specific binding partner through a high-affinity interaction.
- 9. (Original) The method as claimed in claim 8, characterized in that the hapten groups are selected from digoxin, digoxigenin, dinitrophenol and biotin or biotin analogs.

- 10. (Currently Amended) The method as claimed in any of claims 1 to 9 claim 1, characterized in that the hapten groups are applied in a planar fashion to the carrier.
- 11. (Currently Amended) The method as claimed in any of claims 1 to 10 claim 1, characterized in that the hapten groups are applied in a site-specific fashion to the carrier.
- 12. (Currently Amended) The method as claimed in any of claims 1 to 11 claim 1, characterized in that the hapten groups are applied directly to the surface of the carrier.
- 13. (Currently Amended) The method as claimed in any of claims 1 to 12 claim 1, characterized in that the hapten groups are inserted into spacer molecules which are disposed between the carrier surface and the receptors.
- 14. (Currently Amended) The method as claimed in any of claims 1 to 13 claim 1, characterized in that the hapten groups are inserted at one or more positions into the receptors synthesized on the carrier.
- 15. (Currently Amended) The method as claimed in any of claims 1 to 14 claim 1, characterized in that the hapten groups are applied reversibly.

- 16. (Currently Amended) The method as claimed in any of claims 1 to 14 claim 1, characterized in that the hapten groups are applied irreversibly.
- 17. (Original) The use of hapten groups for controlling the synthesis of receptors on a carrier.